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ASSOCIATION OF RADIONUCLIDES WITH STREAMBED SEDIMENTS IN WHITE OAK CREEK WATERSHED

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ABSTRACT

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Radionuclides are found in much higher concentrations on streambed sediment than in the water of White Oak Creek. Selective extraction of sediments demonstrates that 60 Co is immobilized in a nonexchangeable form in the ferromanganese hydrous oxide coatings on the sediments; 90 Sr occurs predominantly in an exchangeable form on clay, iron oxides, and ferromanganese hydrous oxides; 137 Cs occurs in a nonexchangeable and strongly bound form on clays which compose the dominant rock (Conasauga shale) in the watershed. The fine-gravel to coarse-sand size fraction of streambed sediments is the most suitable fraction for radionuclide analysis because of its abundance in the sediment and its high concentration of radionuclides compared to larger and smaller size fractions. A preliminary survey of all major tributaries in White Oak Creek shows that radionuclide analysis of streambed sediments is a very useful technique to locate sources of radioactive contamination.

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TABLE OF CONTENTS

	Page
ABSTRACT	iii
LIST OF TABLES	vii
LIST OF FIGURES	ix
INTRODUCTION	1
EXTRACTION OF METALS AND RADIONUCLIDES AND THEIR RELATIONSHIP TO MINERALOGY	4
Experimental	6
Evaluation of Extractants	6 6 8
Results and Discussion	9
ADSORPTION OF 85SR TO STREAMBED MATERIALS	21
Experimental	22
Lithologic Materials	22 23 24
Results and Discussion	25
OPTIMUM SIZE FOR SEDIMENT SAMPLING	27
Experimental	27
Results and Discussion	28
SURVEY OF WHITE OAK CREEK STREAMBED SEDIMENTS	32
SUMMARY AND CONCLUSIONS	40
REFERENCES	42

LIST OF TABLES

Tab 1	<u>e</u>	Page
1	Extractions used for Fe, Mn, and radionuclides and mechanism of each extractant	5
2	Extraction of Fe, Mn, 60 Co, from coarse sand (3.35 to 0.85 mm) contaminated with 60 Co	10
3	Sequential dissolution of a ferromanganese coating scraped from a shaley limestone cobble from Melton Branch of White Oak Creek. The second set shows that heating the sample to 80 C prior to dissolution has little effect on the extractions	12
4	Extraction of Fe, Mn, 90 Sr, and 137 Cs from coarse sand (3.35 to 0.85 mm) of streambed sediment contaminated with 90 Sr and 137 Cs	15
5	Sequential dissolution of an Fe gel collected at a radioactive groundwater seep in SWDA-4	17
6	Radionuclide and extractable Fe and Mn content of streambed gravel (4.75 to 2.0 mm) segregated by color	19
	Distribution coefficient (K_d) of ^{85}Sr between White Oak Creek water and streambed gravels, sediments, organic matter, and Fe-Mn oxides	26
8	Radionuclide and extractable Fe and Mn of streambed sediment size fractions	31
9	Radionuclide concentrations of streambed sediment samples of White Oak Creek watershed	35

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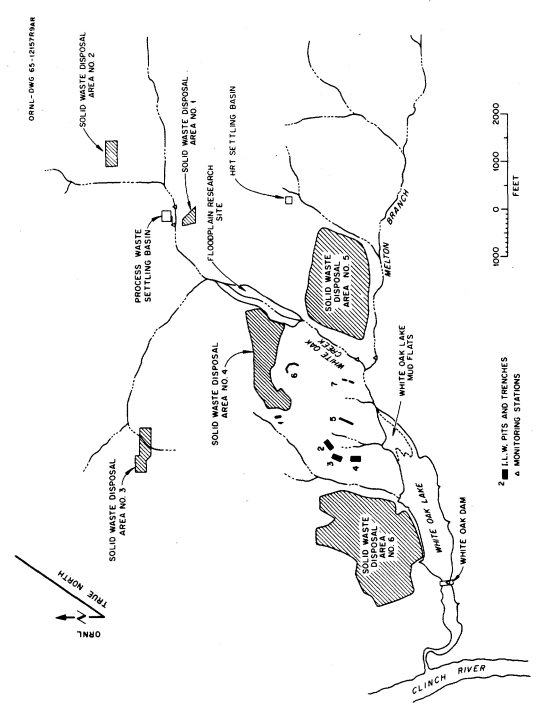
LIST OF FIGURES

Figu	<u>re</u>	Page
1	Location of solid waste disposal areas (SWDA's) and intermediate-level waste (ILW) pits and trenches in White Oak Creek watershed	2
2	The relationship between ^{60}Co and Mn extractabilities by various solutions from a sample of contaminated streambed sediment	11
3	Percent loss of each component from the manganese oxide-coated shale during sequential dissolution	13
4	Percent loss of each component from the Fe sol during sequential dissolution	16
5	The distribution of quartz, ^{137}Cs , and ^{90}Sr , Mn, and Fe extractable by hydroxylamine in size fractions of contaminated streambed sediment	29
6	Location of streambed sediment samples from White Oak Creek watershed. Sample identification numbers correspond to the radiochemical concentrations listed in Table 6	33
7	The distribution of $^{90}\mathrm{Sr}$ concentrations in streambed sediment samples from White Oak Creek watershed	36
8	The distribution of ^{137}Cs concentrations in streambed sediment samples from White Oak Creek watershed	37
9	The distribution of ⁶⁰ Co concentrations in streambed sediment samples from White Oak Creek watershed	38

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INTRODUCTION

Although several general sources of radiochemical contamination entering White Oak Creek (Fig. 1) are known, the estimated releases from these sources fall considerably below the total monitored releases leaving the watershed at White Oak Dam (Stueber et al. 1978). Thus, unknown sources as well as imprecisely located known sources of contamination exist in the watershed. These must be identified before abatement measures can be effectively applied. Analysis of the radiochemical concentrations in stream water and groundwater offers an effective method of locating these sources, but this method has some limitations. Water analysis has a comparatively limited sensitivity, at least when employing sample volumes less than a liter, due to the large dilution of radionuclides by uncontaminated water flowing through the watershed. Sediment analysis offers a considerable increase in sensitivity for radionuclide detection, particularly at point sources of contamination; the laboratory values for distribution coefficients $(K_d = dpm/g \text{ in solid} \div dpm/ml \text{ in water})$ for radionuclides in local groundwater with Conasauga shale, which composes most of the sediment in Melton valley, are high: 120, 70,000 and \geq 100,000 ml/g for 90 Sr, 60 Co, and 137 Cs, respectively. Sediment analysis also provides a short-term history of the surrounding stream water composition, particularly for the more strongly adsorbed radionuclides. In addition, the larger-sized components of stream sediment are transported more slowly by flowing water and, thus, may represent stationary monitors of the radionuclide concentrations in stream waters.



Location of solid waste disposal areas (SWDA's) and intermediate-level waste (ILW) pits and trenches in White Oak Creek watershed. Fig. 1.

This study was undertaken to evaluate the potential use of streambed sediments for locating and monitoring radionuclide contamination in White Oak Creek. Only $^{90}\mathrm{Sr}$, $^{137}\mathrm{Cs}$, and $^{60}\mathrm{Co}$ were examined because these comprise most of the radioactivity, with the exception of tritium, released into White Oak Creek. Each of the radionuclides was examined for its interaction with streambed sediment using several approaches. Contaminated sediment was extracted with water, salt/buffer solutions, and reducing agents to observe the distribution of each radionuclide in various phases: soluble, exchangeable by other cations, or precipitated with ferromanganese hydrous oxides, respectively. The ferromanganese hydrous oxide phase, which coats many of the sediments in White Oak Creek, was examined because of its known importance in the adsorption and immobilization of transition elements (Jenne 1968) including ⁶⁰Co, and because several common manganese minerals contain alkaline earth elements (McKenzie 1977) including 90 Sr. In addition, the distribution of each radionuclide in different size classes of sediment was examined to determine which sizes are most contaminated and, therefore, most suitable for routine monitoring.

Since the initial investigations described above generated some interesting observations, additional and more basic investigations of radionuclide-sediment interactions were initiated. Several hydrous iron oxide precipitates (sols) and isolated ferromanganese hydrous oxide coatings were examined by selective chemical dissolution to elucidate the form and mechanism of radionuclide adsorption and precipitation. Several mechanisms could account for the association of

each radionuclide with these hydrous oxides including coprecipitation, cation exchange by either the oxides themselves, or organic matter inclusions produced by microbial growth or entrapment of particulates from stream water. Particular attention also was focused on the potential interaction of Sr with different types of sediment. Organic detritus in varying stages of decomposition and mineralogically different rocks were examined for their ability to adsorb Sr from stream water spiked with ⁸⁵Sr.

Finally, a preliminary survey was made of the radionuclide content of the fine gravel and coarse sand fractions of streambed sediments from throughout White Oak Creek watershed. Samples were taken from all major streams above and below known or suspected areas of contamination. This information is needed to select areas or sections of streams which will require more intensive sampling to locate sources of contamination entering the watershed.

EXTRACTION OF METALS AND RADIONUCLIDES AND THEIR RELATIONSHIP TO MINERALOGY

Several different extractants were evaluated for their ability to extract iron, manganese, 60 Co, 90 Sr, and 137 Cs from stream sediments. These included cation exchangers, acids, oxidizing agents, chelating agents, and reducing solutions (Table 1). Two different samples were examined: one contaminated with both 90 Sr and 137 Cs and the other contaminated with 60 Co.

Table 1. Extractants used for Fe, Mn, and radionuclides and mechanism of each extractant

	Mechanism ^a					
Extractant	(1)	(2)	(3)	(4)	(5)	Comments
Tapwater						
1 <u>N</u> ammonium acetate pH=7	X			X		
1 <u>N</u> NaC1	Χ					
0.3 <u>M</u> NH4- citrate	X			X		
$^{3\%}$ $^{\text{H}_2\text{O}_2}$ in $^{\text{b}}$ 0.01 $^{\text{M}}$ $^{\text{H}\text{N}\text{O}_3}$		X	х		Х	Reduces Mn (IV) but oxidizes organic matter
0.1 $\underline{\text{M}}$ NH ₂ OH HCL _c in $\overline{\text{0.01}}$ $\underline{\text{M}}$ HNO ₃	Χ	X			X	
2% NH ₂ OH HCL in 0.3 <u>M</u> NH ₄ - citrate, pH 7d	x			Х	X	
0.2 M NH4-oxalate pH Ze	Х	Х		Х		Precipitates Ca, Mg, Sr oxalate
0.2 g/g Na ₂ S ₂ 04 in 0.3 Na citrate, pH 4.8f	х	X		Х	X	
8 <u>N</u> HNO3	X	Х				At high temperature, structural alteration of clay

a(1) Exchangeable (exchange sites on clay, organic matter, Fe-Mn oxides);
 (2) acid (metals, organic matter, carbonates, some silicates);
 (3) oxidizing agent (organic matter);
 (4) chelating agent (Fe and Mn, some organic matter);
 (5) reducing agent (Fe and Mn).

bTaylor et al. 1964.

^CChao 1972.

dWhitney 1975.

eMcKeague and Day 1966.

fCoffin 1963.

In conjunction with this study, sequential dissolutions of a contaminated iron sol and a contaminated manganese coating were carried out to help determine the association of radionuclides with different minerals present in stream gravels.

Experimental

Evaluation of Extractants

Ten-gram duplicates of the ⁹⁰Sr- and ¹³⁷Cs-contaminated sediments were placed in 25- x 150-mm test tubes and extracted twice with 30 ml and once with 20 ml of one of solutions (Table 1) by swirling, waiting one hour, swirling again, and decanting the solution. Combined decantings were filtered by washing through Whatman No. 40 paper to remove the small amount of particulate matter produced during extraction, and were made to 100 ml with water. The extractions of the ⁶⁰Co-contaminated samples were performed on 1.50-g samples with two 15-ml and one 10-ml volumes, filtered as above, and made to 50 ml. All extractions were run in duplicate at both ambient (ca. 25°C) and 95°C temperatures (in a hot water bath). Each extract was analyzed for manganese by oxidation to permanganate (Adams 1965) and for iron by an o-phenanthroline procedure (Jackson 1956).

Sequential Dissolution

A relatively pure Fe sol was obtained from a seep in Solid Waste Disposal Area (SWDA) 4 near well 186A; a ferromanganese hydrous oxide separate was obtained by scraping with a stainless steel spatula the easily flaked-off dark coating on an encrusted shaley limestone cobble

from Melton Branch. Care was taken to avoid cobbles with excessive algal coatings. The iron sample was separated from the liquid phase by centrifugation at 600 RCF for 30 min; the aqueous phase was analyzed after three weeks. The ferromanganese sample was split; one-half was air-dried and the remainder was dried at 80°C. Samples were treated with progressively more destructive reagents and each leachate was then analyzed for Fe, Mn, 60 Co, 90 Sr, and 137 Cs as described later. Weight losses were determined after each step; x-ray diffraction patterns were run after the removal of iron and clay minerals. Reagents used were: ammonium acetate buffered to pH 7 to remove exchangeable cations (Jackson 1956); repeated H_2O_2 treatments followed by extraction with 1 N sodium-acetate at pH 7 to remove organics and MnO_2 (used on Fe sol only); 0.3 \underline{M} ammonium citrate with 2% w/v hydroxylamine hydrochloride adjusted to pH 7.0 with ammonium hydroxide to remove MnO_2 and poorly crystalline iron oxides (Whitney 1975); and Na-dithionite in 0.3 \underline{M} sodium citrate and 0.1 \underline{M} sodium bicarbonate to remove iron oxides (Jackson 1956). These treatments resulted in a residue of disaggregated silicate minerals. This residue (in the case of the iron sample, after separating into sand, silt, and clay fractions) was then treated with hot HNO_3 for 1 hr at 95°C. The final dissolution procedure dissolved clay minerals by fusion with sodium pyrosulphate and leaching with HCl and NaOH (Kiely and Jackson 1964, 1965). The residue was shown to be quartz containing a trace of feldspar, determined by optical examination under a microscope and by X-ray diffraction. Each procedure (except clay dissolution) was

performed twice, followed by three washings with distilled water which were added to the initial supernatant before analysis.

Radiochemical Analyses

A 25-ml aliquot was used for the determination of total radiostrontium (American Public Health Service 1975); the barium carrier addition and its precipitation with sodium chromate were not employed because ¹⁴⁰Ba was not present at detectable levels in these samples. In addition, only one scavaging with the mixed rare-earth carrier was employed because samples were counted after full ingrowth of the $^{90}\mathrm{Y}$ daughter (21 days). The final precipitate (SrCO $_3$) from the radiochemical fractionation was slurried onto a tared 1-in.-diam. aluminum planchet, dried, weighed to calculate carrier recovery (usually between 70 and 80%), and counted on a Beckman Wide Beta II gas-flow proportional counter after 90 Y ingrowth. Counting efficiency was determined using a dilution of standard reference material 4234 (National Bureau of Standards) at various $SrCO_3$ sample thicknesses; typical counting efficiencies were about 42% depending on carrier recovery. The total observed counts, which represent equal contributions from 90 Sr and 90 Y, were halved to report only 90 Sr; this was expressed in dpm/g of oven-dried material.

The extractability of 60 Co and 137 Cs by each solution was determined by computing the difference in radionuclide content of the sediment before and after extraction. Cesium-137 and 60 Co were counted with a Packard Model 16 multichannel analyzer equipped with a well-type NaI (Tl-activated) detector. Counting efficiencies were

determined using single radioisotopic standards (which had previously been compared with NBS standard reference material) prepared in 10-g soil matrices and contained in 25- x 150-mm test tubes. Background counts in the ^{137}Cs channels window were determined by computing the average counts per channel for the five channels on each side of the window and multiplying this figure by the number of channels in the ^{137}Cs window. The background counts in the ^{60}Co channels window were determined by counting empty test tubes because no γ -emitter with decay energies greater than ^{60}Co was present in these samples.

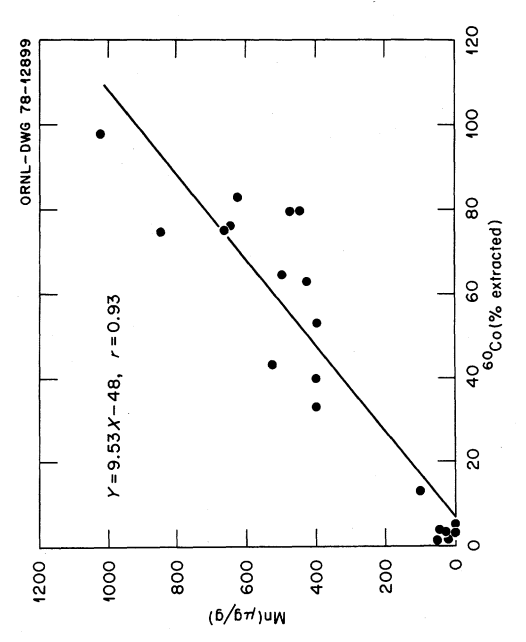
Results and Discussion

Each radionuclide present in sediment was strongly associated with different mineral phases. The extractability of 60 Co in streambed gravel by different reagents strongly suggests its association with the ferromanganese hydrous oxide component in the stream sediment (Table 2). Little 60 Co was water-soluble or exchangeable since tapwater, ammonium acetate, and sodium chloride solutions removed only minor amounts. Other extractants removed varying amounts of 60 Co, Mn, and Fe from the sediment, but the amount of 60 Co extracted was well correlated (r = 0.93) with the amount of Mn extracted (Fig. 2); 60 Co was less well correlated (r = 0.61) with the Fe extracted, because Fe and Mn behaved somewhat independently in their extractabilities (r = 0.65). Sequential dissolution of the scraped-off ferromanganese coating showed that all 60 Co was extracted during the dissolution step which removed 90 to 95% of the Mn but only about 20% of the total Fe in the sample (Table 3 and Fig. 3).

Table 2. Extraction of Fe, Mn, $^{60}\mathrm{Co}$, from coarse sand (3.35 to 0.85 mm) contaminated with $^{60}\mathrm{Co}$

		25°C			95°C	
Extractant	Fe (μg/g)	M n (μg/g)	⁶⁰ Co (% total)	Fe (µg/g)	Mn (μg/g)	⁶⁰ Co (% total) ^a
Tapwater	3	<u><</u> 10	5	2	10	4
1 <u>N</u> ammonium acetate pH = 7	1	20	3	2	45	1
1 N NaCl	2	18	3	6	29	3
0.3 <u>M</u> NH ₄ - citrate	160	90	13	1260	460	80
3% H ₂ O ₂ in 0.01 <u>M</u> HNO ₃	28	410	40	18	400	33
0.1 M NH20H·HC1 in 0.01 M HN03	82	530	44	2830	840	75
2% NH ₂ OH•HCl in 0.3 <u>M</u> NH ₄ - citrate, pH 7	350	430	64	5630	630	84
0.2 M NH ₄ - oxalate pH 2	572	410	54	11750	660	77
0.2 g/g Na ₂ S ₂ 04 in 0.3 Na-citrate pH 4.8	14200	660	77	18760	480	80
8 <u>N</u> HN03	3200	490	65	45270	1020	98

aTotal 60Co was 12,500 dpm/g.



The relationship between $^{60}\mathrm{Co}$ and Mn extractabilities by various solutions from a sample of contaminated streambed sediment. Fig. 2.

Table 3. Sequential dissolution of a ferromanganese coating scraped from a shaley limestone cobble from Melton Branch of White Oak Creek. The second set shows that heating the sample to $80\,^\circ\text{C}$ prior to dissolution has little effect on the extractions

Treatment	Weight ^a loss (mg)	Fe ₂ 0 ₃ (mg)	MnO ₂ (mg)	137 _{Cs} (dpm)	90 _{Sr} (dpm)	60 _{Co}
Original	577.7					
NH ₄ -acetate, pH 7		_b	-	-	113	-
NH4-citrate, 2% NH2OH HCl	(124.1) ^C	5.3	36.4	. .	38	10,000
Na-dithionite		8.1	1.6	- '	-	-
Hot HNO ₃	(29.0)	9.4	-	530	-	-
NaHSO4 fusion	(124.4)	4.9	-	-	~	-
Residue = quartz	300.2	n.d.d	n.d.	-	n.d.	-
Original	577.7					
80 C	(7.0)					
NH4-acetate, pH 7		-	-	- .	109	-
NH ₄ -citrate, 2% NH ₂ OH HCl	(112.3) ^c	5.3	33.5	-	51	10,800
Na-dithionite	l l	7.7	3.8	-	-	-
Hot HNO ₃	(34.4)	10.0	-	580	-	-
NaHSO4 fusion	(125.2)	4.9	-	- ,	-	-
Residue = quartz	298.8	n.d.	n.d.	-	n.d.	-

 $^{^{\}rm a}{\rm Original}$ sample had an air-dry weight of 577.7 mg and contained a small amount of organic matter.

b- = not detected.

CWeight loss after combined treatments.

 $d_{n.d.}$ = not determined.

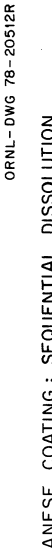
530 dpm

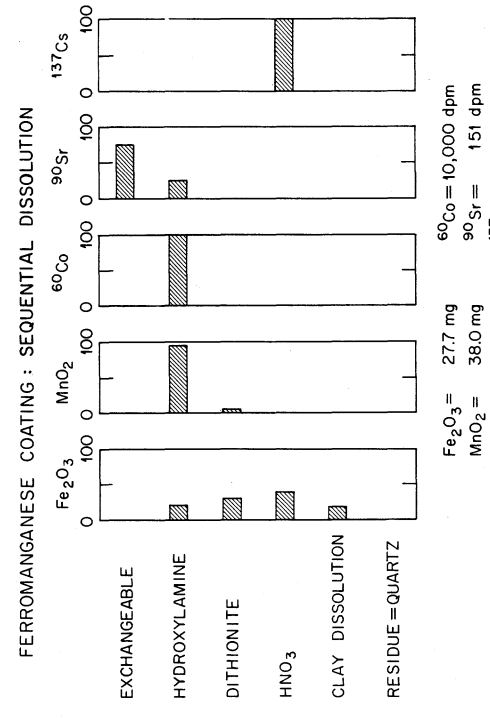
137_{Cs}=

153.4 mg

CLAY=

QUARTZ = 300.2 mg





Percent loss of each component from the manganese oxide-coated shale during sequential dissolution. က် Fig.

Unlike 60 Co, the 90 Sr in streambed sediment was predominantly in an exchangeable form. Although only 4% of the sediment 90 Sr was soluble in tapwater, about 85% was exchangeable since it was removed by hot ammonium acetate or sodium chloride (Table 4). Total 90 Sr is defined as the amount extracted by hot 8 N HNO3. Other extractants were also very effective in removing 90 Sr since they all contain exchangeable cations. Notably, ammonium oxalate and sodium dithionite removed considerably less 90 Sr than ammonium acetate or NaCl; this was likely due to the insolubility of alkaline-earth oxalates and low solubility of Ca and Sr sulfates which would form as dithionite is oxidized. A significant portion, about 15%, of the total 90 Sr was present in a nonexchangeable form.

Sequential dissolution of both the iron sol and the scraped-off ferromanganese coating showed that greater than 70% of the 90 Sr in each material was exchangeable (Tables 3 and 5; Figs. 3 and 4). Hydroxylamine hydrochloride dissolved 90% of the Mn from the scaped-off ferromanganese coating and all of the remaining, nonexchangeable 90 Sr (Table 3 and Fig. 3). The iron sol also lost most of the remaining, nonexchangeable 90 Sr during oxidation by $\mathrm{H_2O_2}$, followed by extraction with sodium acetate and by reduction with hydroxylamine (Table 5 and Fig. 4); no manganese, however, was detected in these extracts. Interestingly, a small amount of 90 Sr was strongly held by the clay phases since hot $\mathrm{HNO_3}$ and sodium pyrosulfate dissolution of the silicate phases was required to remove it (Table 5).

Several other lines of evidence suggest that the ferromanganese coatings contained $^{90}\mathrm{Sr}$ in a nonexchangeable form. In the

Table 4. Extraction of Fe, Mn, $^{90}\rm{Sr}$, and $^{137}\rm{Cs}$ from coarse sand (3.35 to 0.85 mm) of streambed sediment contaminated with $^{90}\rm{Sr}$ and $^{137}\rm{Cs}$

		25°C					95°C			
Extraction	Fe (µg/g)	Mn (µg/g)	90 (dpm,		137 _{Cs} (% total)	Fe (µg/g)	Mn (μg/g)	90 _{Sr} (dpm/g)b	137 _{Cs} (% total)a	
Tapwater	3	<u><</u> 10	37	(2%)	4	1	10	64 (4%)	2	
1 N ammonium acetate pH = 7	6	12	888	(60%)	7	5	. 39	1340 (90%)	10	
1 <u>N</u> NaC1	8	13	789	(53%)	4	6	18	1210 (81%)	4	
0.3 <u>M</u> NH4_ citrate	140	2000	926	(62%)	5	1190	3160	1350 (91%)	6	
3% H ₂ O ₂ in 0.01 <u>M</u> HNO ₃	7	650	316	(21%)	4	6	470	339 (23%)	5	
0.1 M NH2OH•HC1 in 0.01 M HN03	54	2330	749	(50%)	4	891	3740	1260 (85%)	6	
2% NH ₂ OH·HC1 in 0.3 <u>M</u> NH ₄ _ citrate, pH 7	430	2920	1230	(83%)	5	3060	3590	1510 (101%	7	
0.2 <u>M</u> NH4. oxalate pH 2	920	2090	38	(3%)	5	6440	3460	124 (8%)	7	
0.2 g/g Na ₂ S ₂ O4 in 0.3 Na-citrat pH 4.8	e 8220	4270	906	(61%)	6	11,600	3370	841 (56%)	6	
8 <u>N</u> HNO3	5670	1790	1290	(89%)	29	32,900	4920	1490 (100%	49	

aTotal 137Cs was 458 dpm/g.

 $[^]b Total$ $^{90} Sr$ is defined as that extracted by hot 8 $\underline{\text{N}}$ HNO3. Numbers in parentheses are the percent of the total.

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Fe203 SOL: SEQUENTIAL DISSOLUTION

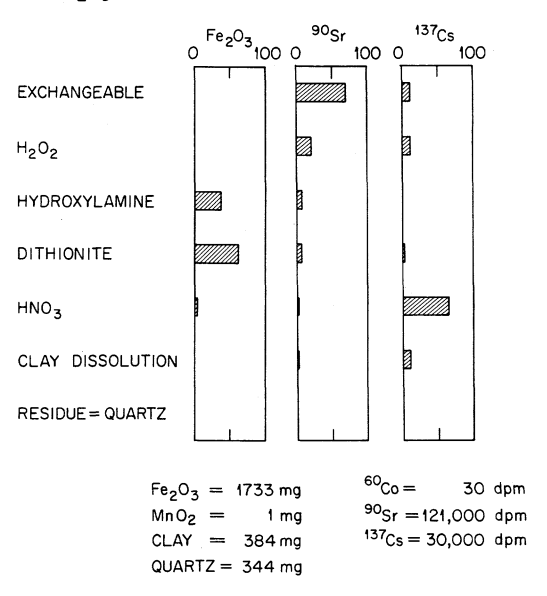


Fig. 4. Percent loss of each component from the Fe sol during sequential dissolution.

Table 5. Sequential dissolution of an Fe sol collected at a radioactive groundwater seep in SWDA-4

Treatment	Weight ^a loss (mg)	Fe ₂ 0 ₃ (mg)	MnO ₂ (mg)	137 _{Cs} (dpm)		60 _{Co} (dpm)
Water	n.d.b	_c	-	8.0d	100 ^d	3.3d
1 N NH4 acetate pH 4.5	n.d.	0.3	1.3	3310	84,000	30
H ₂ 0 ₂	n.d.	1.6	-	3150	24,000	-
NH ₄ -citrate, 2% NH ₂ OH HC1	n.d.	656	-	440	6,000	-
Na-dithionite	n.d.	1066	-	850	6,200	-
Hot HNO $_3$ > 63 μ 63 to 2 μ $_<$ 2 μ	(9.5)e (51.6)e (39.2)	0.5 3.1 3.0	- - -	507 5360 13900	- 62 48	- - -
NaHSO4 fusion > 63 63 to 2 μ < 2 μ	(22.9) (95.2) ^e (167.1)	0.1 0.9 0.9	0.1	- - 2760	- 24 188	- - -
Residue = quartz > 63 63 to 2 μ < 2 μ	142.1 174.2 ^e 27.3	n.d. n.d. n.d.	n.d. n.d. n.d.	- - -	- - -	- - -

 $^{^{\}mathrm{a}}$ The original sample had an oven-dry weight of 3.30 g and contained some organic matter.

bn.d. = not determined.

c- = not detected.

dDpm/ml.

^eCentrifuge tube broke. Extractions carried out on broken tube plus samples. Weights estimated by comparison with another sample.

non-sequential extractions (Table 4), although about 85% of 90 Sr was exchangeable, all 90 Sr was removed by extraction with hydroxylamine in ammonium acetate; this extractant removed most of the Mn but only 10% of the Fe extracted by hot 8 M HNO3 (Table 4). Secondly, when a ferromanganese-coated sediment sample was segregated by color into a light and dark fraction, the dark-colored gravel contained 27% more 90 Sr than the lighter-colored gravel (Table 6). The dark-colored gravel contained five times as much Mn and twice as much Fe extracted by hydroxylamine than the lighter colored gravel. These separate observations indicate that, although 70 to 85% of the 90 Sr was held in an exchangeable form by streambed gravels and their ferromanganese coatings, most of the remaining, nonexchangeable 90 Sr was associated with ferromanganese oxides. Trace amounts were also held in a nonexchangeable form by clay minerals.

Cesium-137, unlike either 90 Sr or 60 Co, could not be extracted in significant amounts (Tables 2 and 4) by any solution other than 8 $\underline{\text{M}}$ HNO $_3$ which causes structural changes in crystalline minerals. This lack of extractability of 137 Cs illustrates its strong and specific adsorption by illitic clays (Tamura and Jacobs 1960, Tamura 1963) which compose the parent rock, Conasauga shale, of which this streambed gravel was composed. Dark-colored gravel from a split sample contained approximately five times as much 137 Cs and Mn as did the light-colored gravel (Table 6). Progressive dissolution of the scraped-off ferromanganese coating showed that all the 137 Cs was associated with illite since the 137 Cs was not dissolved until treatment with 8 $\underline{\text{M}}$ HNO $_3$ (Table 3 and Fig. 3). Dissolution of the

Table 6. Radionuclide and extractable Fe and Mn content of streambed gravel (4.75 to 2.0 mm) segregated by color

	Light-colored gravel	Dark-colored gravel	
137 _{Cs} (dpm/g) ^a	54	266	
90 _{Sr} (dpm/g)b	1025	1300	
60 _{Co} (dpm/g)a	6.0	20.0	
Mn (μg/g) ^b	910	4630	
Fe (μg/g) ^b	1220	2940	

aTotal content based on triplicate 10-g samples.

 $[^]b Extractable$ in 2% hydroxylamine hydrochloride/0.3 $\underline{\text{M}}$ ammonium citrate, pH 7, based on triplicate 10-g samples.

iron sol was less straightforward (Table 5 and Fig. 4). Although most of the ^{137}Cs was released by 8 M HNO_3 or by treatment to remove the layer silicates, significant amounts were present in the water associated with the seep or removed by ammonium acetate or H_2O_2 extraction. The high concentration of ^{137}Cs in the water likely indicates that the adsorption capacity of the small amount of clay in this sample may have been exceeded. The calculated K_d of ^{137}Cs for this sample and its associated water was about 10,000.

These studies have shown which mineral phases are responsible for the adsorption of each radionuclide by streambed sediment. They also give some indication of how each radionuclide is likely to move in the stream. Because ⁶⁰Co was found to be strongly associated with the ferromanganese oxide phase, it is likely to be solubilized by stream water as sediment moves from well-aerated to reducing sections of a particular stream. The ferromanganese hydrous oxide coatings on streambed sediments undergo deposition or dissolution, depending on the Eh of the stream water (Jenne 1968). Cobalt-60 is, therefore, likely to move in streams as Mn does.

Cesium-137 was also restricted almost entirely to a single phase. Even the ferromanganese-coated sample had all of its 137 Cs associated with the layer silicate phase; most of 137 Cs in the iron sol was also associated with the layer silicate phase. Cesium-137 content was also five times greater in dark-colored streambed gravel than in the portion of the same sample which was lighter-colored (Table 6). These results suggest that 137 Cs-clay particles may be incorporated into growing ferromanganese coatings. The ferromanganese coating contained less

than 10% Mn plus Fe; the remainder was original rock, detrital minerals, and, possibly, organic matter. Examination of an apparently homogeneous ferromanganese coating with an electron microprobe (10- μ beam diam.) showed a maximum of 42% MnO₂ and 14% Fe₂O₃, with minor amounts of Si, Al, K, Ca, Ni, and Cu indicating the presence of silicate inclusions as well as other metals.

Strontium-90 occurred primarily as an exchangeable cation in streambed sediment, although it is uncertain which of the three dominant phases (clay, δ -MnO $_2$ or Fe-sol) was most important. Iron and Mn hydrous oxides have a significant capacity to adsorb cations (Kinniburgh et al. 1976, Loganathan et al. 1977) and several common manganese minerals contain alkaline earth elements (McKenzie 1977). Although some 90 Sr is likely in a nonexchangeable form, most 90 Sr is likely partitioned between the stream water and sediment by a cation exchange mechanism probably maintained by the level of dissolved Ca in stream-water.

ADSORPTION OF 85sr TO STREAMBED MATERIALS

As noted above, several different materials have cation exchange sites available for strontium. In order to estimate the relative importance of each of these materials in radionuclide contamination, distribution coefficients for trace levels of strontium were determined for bedrock materials, various kinds of organic matter, and iron and manganese hydrous oxides. Samples of streambed gravel were collected on March 20 and 21, 1978, as part of a preliminary survey of White Oak Creek sediment for radionuclide contamination. From this collection,

several samples were selected that had differing lithologies: brown shale, dark shale with ferromanganese coating, light-colored chert, limestone, chert-limestone-shale mixture, and dark chert with ferromanganese coating. Samples of organic material were collected from White Oak Creek upstream (northeast) of the ORNL plant site.

Experimental

Lithologic Materials

One milliliter of carrier-free ⁸⁵Sr (138,000 dpm) was added to 25 ml of freshly collected creek water (pH 7.3, e.c. 96 µmhos/cm) containing 10.0 g of oven-dried gravel-sand in a 50-ml polypropylene centrifuge tube. The tube was shaken lengthwise for 18 hr at room temperature and centrifuged at 650 RCF for 10 min on an IEC model PR-2 refrigerated centrifuge. A 10-ml aliquot of the supernatant was transferred to a 30-ml polycarbonate "Oak Ridge"-type centrifuge tube and centrifuged for 15 min at 36,000 RCF with a Beckman L-2 ultracentrifuge (equipped with a No. 30 head) to remove suspended material not removed during the initial centrifugation at lower speed. A 5-ml aliquot of the supernatant was transferred to a 25- \times 150-mm test tube and counted on a Packard Model 16 multichannel analyzer using 30,000 dpm of 85 Sr in 5 ml of water as a standard; activity of the standard was determined by the Analytical Chemistry Division. All observations were corrected for radioactive decay from the time of calibration. Background counts in the $^{85}\mathrm{Sr}$ channels window were obtained with empty test tubes. The amount of $^{85}\mathrm{Sr}$ adsorbed by a sample was determined by calculating the difference in the concentration in the supernatant of a sample and that of a control containing no gravel. The concentration of $^{85}\mathrm{Sr}$ in the creek water was determined directly from the counts of the 5 ml of supernatant. The average K_d from duplicate determinations was calculated by the ratio of dpm/g of solids to dpm/ml of supernatant.

Organic Materials

Samples of organic materials common to streambeds were collected from White Oak Creek, upstream (northeast) of the ORNL plant site. These included a sound wood branch, a somewhat darkened/rotted wood branch, a well-rotted wood branch, partially discolored deciduous leaves, dark-colored/well-rotted leaves, and a green algal slime. The material was oven-dried (100°C, overnight) and ground in a Wiley mill to pass a 2-mm sieve. Duplicate samples (2.50 g) of each type were shaken with 35 ml of freshly collected creek water (pH 7.7, electrical conductivity of 196 μ mhos/cm) to which was added 109,000 dpm of 85 Sr in 1 ml of water. The average K_d for each material was calculated after counting supernatants, as described above. A second K_d determination for these same organic materials was performed after twice washing the samples with water. Duplicate 1-g samples were weighed into tared 50-ml polypropylene centrifuge tubes. Forty milliliters of freshly collected creek water (pH 7.7, electrical conductivity of 196 mhos/cm) were added and the tubes capped and shaken lengthwise for 48 hr. The tubes were centrifuged for 15 min at 650 RCF on the IEC Model PR-2 centrifuge and the supernatants discarded; a second wash was carried out with 30 ml of creek water with shaking for 4 hr, centrifuging, and decanting as before. The tubes were weighed to determine the amount of adsorbed water and additional creek water added to attain 30 ml per tube. One milliliter of water containing 97,000 dpm of 85 Sr was added and the contents shaken lengthwise for 16 hr. The K_d was then calculated as described above.

Synthetic Iron and Manganese Hydrous Oxide

Artificial $^{\delta}$ -MnO $_2$ and iron hydrous oxides were prepared by the oxidation of a MnCl $_2$ -NaOH mixture at pH 8 to 12 (similar to the method of Jeffries and Stumm 1976) and by neutralization of Fe(NO $_3$) $_3$ by NaOH (Kinniburgh et al. 1975) to pH 7, respectively. After formation of the hydrous oxides, excess salts were removed by repeated washings with distilled water and centrifugings at 600 RCF for 20 min until the hydrous oxide began to destabilize. The hydrous oxide was then air-dried. Twenty milliliters of White Oak Creek water (pH 7.7, electrical conductivity of 196 μ mhos/cm) were added to 50 to 100 mg of air-dried gel; 85 Sr (86,000 dpm) was added to each sample. Samples were then shaken continuously for 24 hr and analyzed as described above.

A second set of hydrous oxides was prepared by the oxidation or neutralization of aliquots containing a known amount of manganese or iron (50 to 100 mg). These samples were treated as above except that the air drying step was omitted to minimize structural changes due to drying. The amounts of manganese or iron hydrous oxide in these samples were checked for loss by filtering blank precipitates through tared fritted-glass crucibles. Oven-dried (110°C) sols weighed within

5% of the amount of MnO_2 or Fe_2O_3 in the aliquot, indicating that loss was minimal.

Results and Discussion

Table 7 demonstrates that Conasauga shale had a six- to sevenfold greater $\mathbf{K_d}$ for $^{85}\mathrm{Sr}$ than chert and a tenfold greater $\mathbf{K_d}$ than limestone. Notably a ferromanganese hydrous oxide-coated shale (dark-colored) exhibited a 12% greater K_d for $^{85}\mathrm{Sr}$ than an uncoated (light-colored) shale. Rotted wood and leaves exhibited an affinity for $^{85}\mathrm{Sr}$ similar to Conasauga shale (Table 7). Oven-drying of wood and leaves leads to a release of some soluble organic matter (Jenkinson 1966), and it was suspected that this might increase the ionic strength of the stream water that was used for the ${\rm K}_{\rm d}$ determination, and hence, lead to an unrealistically low K_d . The equilibrium supernatant from the unwashed, oven-dried organic materials appeared highly colored; twice prewashing these organic materials with creek water removed most of the color and in some cases, the observed K_d 's rose considerably. The K_d 's determined with the prewashed organic materials likely reflect their affinities for $^{90}\mathrm{Sr}$ in the unperturbed (non-oven-dried) state.

Several K_d's were determined for natural and synthetic iron hydrous oxides and δ -MnO₂. These K_d's were quite variable and probably were related to the degree of crystallinity of the specimens. Burns (1976) suggested that the adsorbtion of cations should decrease as the amount of birnessite (crystalline δ -MnO₂) increases. In addition, the adsorption capacity of both iron sols and δ -MnO₂

Table 7. Distribution coefficient (K_d) of $^{85}\mathrm{Sr}$ between White Oak Creek water and streambed gravels, sediments, organic matter and Fe-Mn oxides

Gravels	Sample number	K _d ^a	
Light-colored shale Dark-colored shale	9-7 9-8	129 144	b b
Light-colored chert	9-14	20	b
Mixed chert and limestone	9-18	23	b
Mixed chert and limestone Limestone	9-19 9-21	8 6	b b
Organic material	K _d (unwashed) ^d	K _d (washed)	
Wood, light-colored (sound)	49	60	С
Wood, dark-colored (rotted)	102	144	С
Wood, very dark (rotted) Deciduous leaves: dark,	78	155	С
slightly decomposed	127	392	С
Deciduous leaves: very dark	 -	***	J
slightly decomposed	101	160	С
Algal slime	31	55	С
Fe-Mn oxides	K _d		
Fe ₂ 0 ₃ -sol (natural)	550		С
Fe ₂ 0 ₃ -sol (synthetic)	150		C
δ-MnO2 natural	100	to > 1000	c
δ -Mn $0\frac{1}{2}$ synthetic	100	to > 1000	С

 $^{{}^{}a}K_{d} = \frac{dpm/g \ solid}{dpm/ml \ creek \ water}$.

^bCreek water of pH 7.3, ϵ = 96 μ mhos/cm.

^CCreek water of pH 7.7, ϵ = 196 μ mhos/cm.

 $^{^{}d}\text{Prewashed}$ samples rinsed twice with this creek water before K_{d} determinations (see text).

changes most rapidly in the pH range of 6 to 9 (Kinniburgh et al. 1975, Loganathan et al. 1977) which includes the pH (7-8) of the stream water used for these K_{d} determinations. Thus, small changes in pH due to aging of the gel could lead to marked changes in the measured K_{d} 's.

OPTIMUM SIZE FOR SEDIMENT SAMPLING

Iron and manganese concentrations are not uniformly distributed with respect to the size of sediments (Whitney 1975). Because of mineralogical differences between different size fractions, it is expected that the radionuclides will not be uniformly distributed either. Furthermore, it would be useful to sample a size that is easily reproducible, one that can be done in the field, and one that is present in large amounts at most sites. For these reasons, the distribution of radionuclides with respect to size fraction was determined.

Experimental

On March 1, 1978, a large (approximately 3.5-kg) sample of sediment was collected to a 4-cm depth from the channel of the tributary of White Oak Creek that runs south of SWDA-4. The sample was sieved under a flow of tapwater into seven size fractions; clay and silt fractions were isolated by gravity or centrifugal sedimentation (Jackson 1956). All gravitational and centrifugal size isolates were washed three to five times by resuspending in tapwater until the final

supernatant appeared clear; all supernatants were combined and used for the isolation of the next smaller size fraction. Each fraction was oven-dried (105°C, overnight) and weighed.

Each size fraction was analyzed for $^{137}\mathrm{Cs}$ content before and after extraction with 2% hydroxylamine hydrochloride/0.3 M ammonium citrate, pH 7.0, as described previously. The $^{90}\mathrm{Sr}$, Mn, and Fe contents were also determined on these extracts as described previously. The 60 Co content of each fraction (except the coarse-silt and all clay fractions which were expended) was also determined as described previously. The quartz contents were estimated by comparing quartz X-ray diffraction peak heights (100) in each size fraction with that in artificial quartz-illite mixtures. Standards and samples were ground to pass a 325-mesh sieve and mounted in reversible glass-backed aluminum frames. Each standard was run three times to prepare a calibration curve; unknowns were also run three times and the average peak height was used to estimate the quartz percentage. A Philip's X-ray diffractometer with a Ni-filtered Cu-K radiation source operating at 40 kv and 20 ma with a scanning speed of 1°/min was used for these studies.

Results and Discussion

The distribution of ^{137}Cs , ^{90}Sr , Fe, and Mn in size-fractionated sediment further illustrates associations between radionuclides and mineral phases (Fig. 5). Both ^{90}Sr and ^{137}Cs concentrations were bimodal in this sediment, with maxima in the fine gravel to coarse sand (-2 to +2 ϕ) and again in the medium-silt to clay

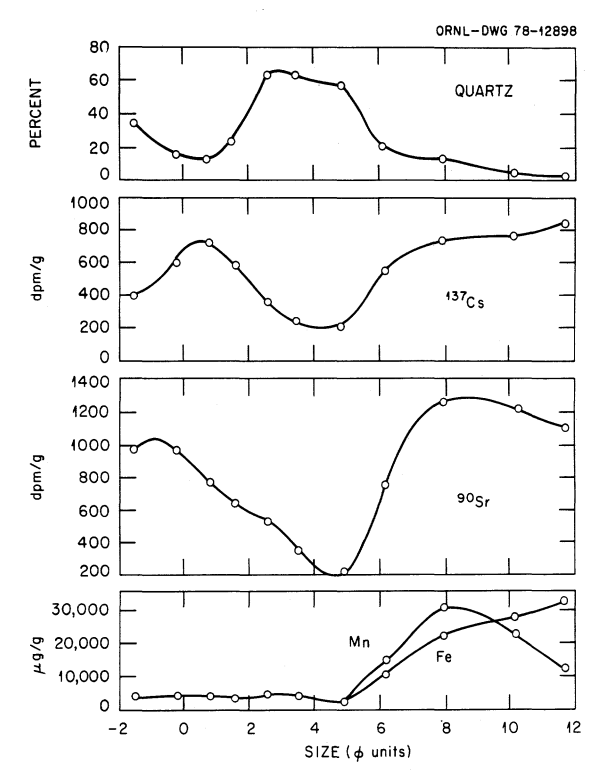


Fig. 5. The distribution of quartz, ^{137}Cs , and ^{90}Sr , Mn, and Fe extractable by hydroxylamine in size fractions of contaminated streambed sediment.

fractions (7 to 9 ϕ). Cobalt-60, although present at only low levels in this sample, also exhibited a maximum in the coarse-sand fraction (Table 8) and appeared to be increasing again from the coarse- to medium-silt-sized fractions; unfortunately, the finer-sized fractions were not available for ⁶⁰Co analyses. Since the size isolates larger than medium sand represented greater than 95% of whole sample weight (Table 8), most of the total radioactivity was associated with these larger fractions; such larger-sized fractions would also have a much lower potential than the finer sizes for transport in the stream as suspended solids. Thus, the coarser fractions will have a longer residence time in a given stream section and their radionuclide content likely represents prolonged contact with stream water within that stream section. Ideally, a stationary fraction, such as exposed bedrock, could be chosen but this is rarely possible. Thus, the fine-gravel to coarse-sand fraction appears to be the best size fraction for monitoring radionuclide contamination by 60 Co, 90 Sr, and ¹³⁷Cs of streambed sediments because of its abundance in the sediment, its ease in isolating and sampling, and its high concentrations of the radionuclides of interest. In addition, gravels/sands with ferromanganese hydrous oxide coatings represent those sediments which have been deposited for some time and, hence, would have more abundant sites for radionuclide retention. These coated sediments generally occur in riffles in the streams.

The bimodal distribution of each radionuclide was strongly related to the mineralogical composition of each size fraction. Quartz, with its comparatively inert surface, would not be expected to adsorb

Radionuclide and extractable Fe and Mn of streambed sediment size fractions Table 8.

				Size fraction	action					
Name	Range (µ)	Mean size (¢)	Weight (g)	Mna (µg/g)	Fea (µg/g)	90 _{Sr} a (dpm/g)	137 _{Cs} b (dpm/g)	60 _{Co} b (g/mdb)	Quartz (%)	Notes
Coarse gravel Fine gravel Very coarse sand Coarse sand Medium sand Fine sand Very fine sand Coarse silt Medium silt Fine silt Coarse clay Medium clay	40,000-4750 4,750-2000 2,000-850 850-500 500-212 212-125 125-88 88-20 20-12 12-2 20-12 20-12 20-12 20-12 20-12	-3.79 -1.63 -0.39 +0.62 1.62 2.62 3.26 4.58 6.01 7.68 9.97	2200.0 741.8 325.1 100.6 58.2 19.1 17.8 10.9 31.2 19.4 6.2 1.6	2700 4130 4770 4170 3760 4920 4920 4310 2140 15700 31900 23800 13000 4210	2110 4070 4220 3500 3600 5270 3970 2240 11300 23200 23200 23200 23200 27400	539 996 942 776 653 653 547 366 216 764 1228 1123 408	n.d.c 388 583 709 568 348 240 214 550 732 758 846 675	n. d. 13 26 30 29 26 18 n. d. n. d. n. d.	n.d. 33 16 13 24 63 63 63 63 63 - h	

 $^{\rm a}$ Extracted in 2% NH $_{\rm 2}$ OH·HCl/0.3 $\underline{\rm M}$ ammonium citrate, pH 7.

^bTotal.

cn.d. = not determined.

driplicate 10-g samples used for extraction and counting.

eTriplicate 5-g samples.

fDuplicate 5-g samples.

^gSingle 5-g sample. h = not detected.

ⁱSingle 1-g sample.

jSettled solids 24 hr after flocculation with 50 ml of $1\underline{M}$ CaCl₂ in 8 liter of combined rinse water from previous size isolate.

^kSingle 0.95-g sample.

radionuclides. Notably its concentration reached a maximum in the fine sand to coarse silt fractions exactly where ⁹⁰Sr and ¹³⁷Cs concentrations on the sediment attained minima. The larger-sized fractions were dominated by shale fragments, i.e., aggregated clay and other minerals, with considerable ferromanganese hydrous oxide coatings; quartz exists as sand- and silt-sized grains in the shale fragments of the coarser sizes. Quartz is the dominant mineral in the medium-sand to coarse-silt fraction, while clay minerals dominate in the smaller fractions. This large increase in ferromanganese coatings in the clay-sized fractions was probably due to the large increase in surface area per unit weight with decreasing particle size.

The distribution of radionuclides and mineralogical composition with sediment size will differ when other rock types dominate the sediment. However, Conasauga shale is the dominant parent rock in Melton Valley where, presumably, most radionuclide contamination arises. Other types of rocks including chert, dolomite, and limestone enter White Oak Creek in Bethal Valley and are transported into Melton Valley by the creek.

SURVEY OF WHITE OAK CREEK STREAMBED SEDIMENTS

Samples of streambed sediment were collected from all major streams and tributaries in the White Oak Creek drainage basin on several dates in March and April 1978. Figure 1 shows the location of the solid waste disposal areas (SWDA) and intermediate-level liquid waste (ILLW) pits and trenches within the White Oak Creek basin; Fig. 6 shows the location of sampling sites with respect to these references

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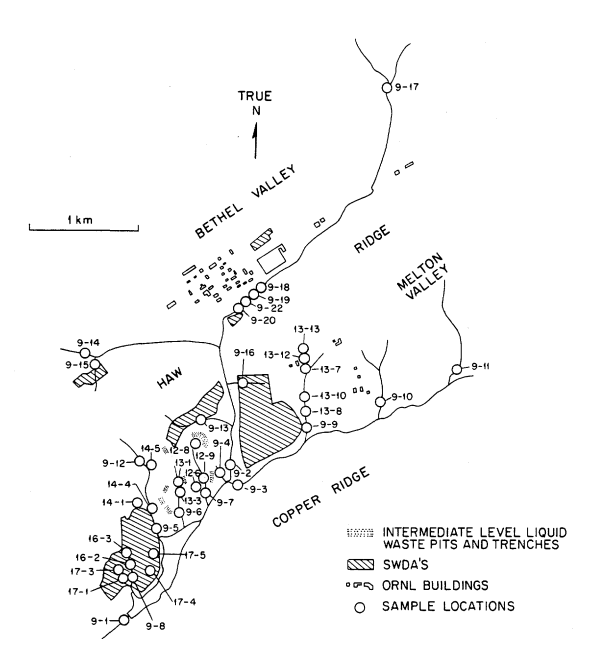


Fig. 6. Location of streambed sediment samples from White Oak Creek watershed. Sample identification numbers correspond to the radiochemical concentrations listed in Table 6.

and ORNL buildings. Samples were collected from sections of streams where ferromanganese oxide-coated gravel deposits appeared to dominate the surface of the streambed sediment; generally, these areas are narrower and more rapidly flowing sections of a given stream. Samples were collected to a depth of approximately 4 cm and placed in polyethylene seal-lock bags. The material was sieved under flowing tapwater and the fine-gravel and coarse-sand fraction (3.35 to 0.85 mm) isolated and oven-dried (105°C, overnight). The 60 Co and 137 Cs content of a 10-g sample was determined as described previously. The exchangeable 90 Sr content was determined using the hot 1 N ammonium acetate, pH 7, extracting procedure described previously.

The observed radionuclide content of each sample is listed in Table 9. To summarize the distribution in the watershed, a graphical representation of each radionuclide's distribution is presented in Figs. 7 (90 Sr), 8 (137 Cs), and 9 (60 Co).

Several generalizations concerning the origin and distribution of each radionuclide can be made. First, the headwater sediments of each stream were at or near background levels of radioactivity of each radionuclide. Strontium-90 appeared to have several major origins including SWDA-4, SWDA-5, ORNL plant effluents, the stream draining the western half of SWDA-6, ILLW trench 6, and ILLW pit 1 (samples 9-13, 13-10, 9-19, 16-2, 12-8, and 14-5, respectively). Cesium-137 seemed to be distributed similarly to ⁹⁰Sr but did not appear in SWDA-6 or ILLW pit 1 drainage; additional low contamination (41 and 26 dpm/g) was observed in the sediment of streams draining SWDA-3 (sample 9-15) and the high flux isotope reactor complex (sample 9-10), respectively.

Table 9. Radionuclide concentrations of streambed sediment samples of White Oak Creek watershed

Sample No.	Total 137 _{Cs} (dpm/g)	Total ⁶⁰ Co (dpm/g)	Exchangeable ⁹⁰ Sr (dpm/g)
9-1	969	139	17.1
9-2	3350	203	22.2
9-3 9-4	57 _a	119	28.5 0.1
9 - 4 9 - 5	11	10,220 541	43.0
9-6	_	1,053	1.4
9-7	25	170	3.3
9-8		3	104.3
9-9	3331	8	61.0
9-10	26	1199	0.1
9-11	-	4	-
9-12	246	2	0.2
9-13 9-14	246	21	1365.1 1.3
9-14 9-15	- 41	3	4.7
9-16	- -	4	0.5
9-17	-		0.7
9-18	9	3	-
9-19	164	47	107.4
9-20	1108	42	13.2
9-22	3409	554 225	17.6 0.6
12 -6 12-8	8 136	335 38	148.0
12-9	62	370	5.1
13-3	5	440	3.3
13-7	95	180	4.2
13-8	893	3	8.3
13-10	1570	7	52.0
13-12	9	104	2.5
13-13	-	3 1	1.0 0.9
14-1 14-4	- 9	25	30.5
14-5	4	<u> </u>	54.0
16-2	-	3	87.5
16-3	-	1	0.9
17-1	-	1	0.3
17-3	-	1	0.3
17-4	-	25 5 3 1 1 2 1	0.9
17-5	-	· 1	0.8

 a_- = not detected.

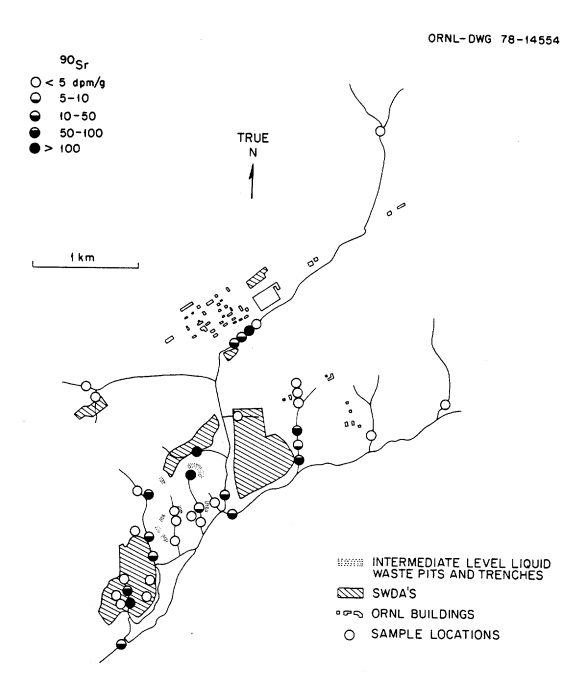


Fig. 7. The distribution of $^{90}\mathrm{Sr}$ concentrations in streambed sediment samples from White Oak Creek watershed.

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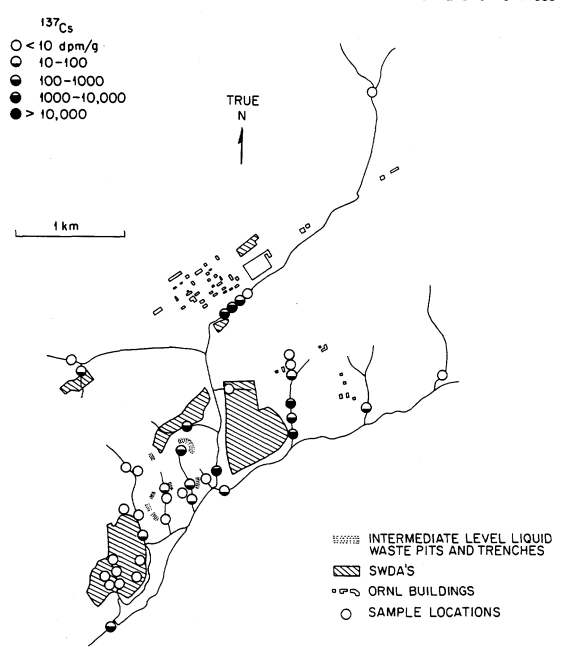


Fig. 8. The distribution of ^{137}Cs concentrations in streambed sediment samples from White Oak Creek watershed.

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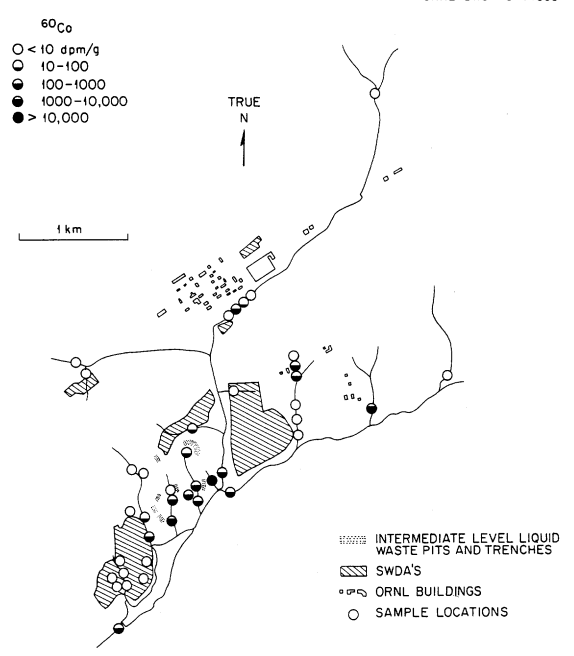


Fig. 9. The distribution of $^{60}\mathrm{Co}$ concentrations in streambed sediment samples from White Oak Creek watershed.

Rather significant 137 Cs concentrations (\geq 3000 dpm/g) were observed at the mouth of the stream east of SWDA-5 (sample 9-9), in White Oak Creek just below monitoring station 3 (sample 9-2), and near points of entry of ORNL plant effluents into White Oak Creek (sample 9-22).

Cobalt-60 exhibited a distribution (Fig. 9) quite different from either 90 Sr or 137 Cs. Major sources appear to be ILLW trenches and pits, particularly the well-known seep east of ILLW trench 7 (Means et al. 1978), plant effluent from the main ORNL complex, leakage near the HRI Settling Basin area, and the high flux isotope reactor (HFIR) area (samples 9-4, 9-22, 13-12, and 9-10, respectively). In general, the low concentrations of 60 Co in the SWDA drainage show that only small amounts, relative to 90 Sr and 137 Cs, originate from the SWDA's. Although the ILLW pits and trenches are repositories for all three of these radionuclides, only 60 Co appears to be moving from them. This suggests that the treatment of the intermediate-level liquid waste (high Na and pH) may have led to immobilization of 90 Sr and 137 Cs in the ILLW pits and trenches. A similar treatment of low-level solid waste or of the soil in which it is to be buried might well lead to better retention of 90 Sr and 137 Cs.

More extensive sampling of streambed sediment is being performed to locate points of entry of radionuclide contamination. Particular attention is being focused on areas where the preliminary samplings indicated large and unknown sources of contamination upstream. Even if radionuclide contamination of stream-water were to be stopped, the sediments themselves pose as a source of contamination for some as yet unestimated time. To estimate how long streambed sediments will remain

a source of contamination, samples of highly contaminated fine gravel, contained in nylon-mesh bags, will be placed in uncontaminated sections (headwaters) of White Oak Creek. By removing and analyzing these samples at various times, estimates of the in situ rates of radionuclide transfer from sediment to stream water under local conditions can be calculated. In addition, a similar experiment, placing uncontaminated gravel in sections of contaminated streams with periodic sampling, should yield a good estimate of the rate of transfer of radionuclides from water to sediment as well as reveal whether contamination is continuing or a result of earlier releases. These two rates should provide enough knowledge to estimate how long streambeds would remain contaminated by each radionuclide as well as to estimate their rate of transport in the creek. Studies of the concomitant rates of ferromanganese hydrous oxide deposition on streambed sediment should further clarify their role in the adsorption and immobilization of 60 Co, 137 Cs, and 90 Sr in White Oak Creek.

SUMMARY AND CONCLUSIONS

This study was initiated to determine how streambed sediments could be used to locate radionuclide contamination in White Oak Creek watershed. Several conclusions were reached:

(1) The three radionuclides, ⁶⁰Co, ⁹⁰Sr, and ¹³⁷Cs, are retained on streambed sediment by different mechanisms. Cobalt-60 is precipitated with manganese in hydrous oxide coatings on minerals or rock fragments. Strontium-90 occurs primarily as an exchangeable cation on clays, organic matter, iron oxides, and manganese oxides;

small amounts of 90 Sr also occur in a nonexchangeable form in manganese oxides. Cesium-137 is held irreversibly by illitic minerals in the sediments.

- (2) Fine gravel to coarse sand is the best size fraction of Conasauga shale-derived sediments for sampling radionuclide contamination. Medium to very fine sand contains more quartz than larger or smaller fractions. Clay and silt size fractions, although they contain radionuclides, are not useful for sampling because they comprise only a small percentage of the stream bottom sediment and because they cannot be readily isolated by sieving when sampled in the field.
- (3) White Oak Creek watershed contains several previously unreported sources of contamination such as 90 Sr from SWDA-6. A much more comprehensive sampling of White Oak Creek sediments is underway to locate with more precision all sources of contamination entering the watershed.

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